

Food and Drug Administration 10903 New Hampshire Avenue Document Control Center – WO66-G609 Silver Spring, MD 20993-0002

June 11, 2015

Nuvasive, Incorporated Martin A. Yahiro, M.D. Director, Medical Affairs 7475 Lusk Boulevard San Diego, California 92121

Re: K150362

Trade/Device Name: NuVasive® CoRoent® Small Interbody System

Regulation Number: 21 CFR 888.3080

Regulation Name: Intervertebral body fusion device

Regulatory Class: Class II Product Code: ODP

Dated: May 14, 2015 Received: May 15, 2015

#### Dear Dr. Yahiro:

We have reviewed your Section 510(k) premarket notification of intent to market the device referenced above and have determined the device is substantially equivalent (for the indications for use stated in the enclosure) to legally marketed predicate devices marketed in interstate commerce prior to May 28, 1976, the enactment date of the Medical Device Amendments, or to devices that have been reclassified in accordance with the provisions of the Federal Food, Drug, and Cosmetic Act (Act) that do not require approval of a premarket approval application (PMA). You may, therefore, market the device, subject to the general controls provisions of the Act. The general controls provisions of the Act include requirements for annual registration, listing of devices, good manufacturing practice, labeling, and prohibitions against misbranding and adulteration. Please note: CDRH does not evaluate information related to contract liability warranties. We remind you, however, that device labeling must be truthful and not misleading.

If your device is classified (see above) into either class II (Special Controls) or class III (PMA), it may be subject to additional controls. Existing major regulations affecting your device can be found in the Code of Federal Regulations, Title 21, Parts 800 to 898. In addition, FDA may publish further announcements concerning your device in the <u>Federal Register</u>.

Please be advised that FDA's issuance of a substantial equivalence determination does not mean that FDA has made a determination that your device complies with other requirements of the Act or any Federal statutes and regulations administered by other Federal agencies. You must comply with all the Act's requirements, including, but not limited to: registration and listing (21 CFR Part 807); labeling (21 CFR Part 801); medical device reporting (reporting of medical device-related adverse events) (21 CFR 803); good manufacturing practice requirements as set

forth in the quality systems (QS) regulation (21 CFR Part 820); and if applicable, the electronic product radiation control provisions (Sections 531-542 of the Act); 21 CFR 1000-1050.

If you desire specific advice for your device on our labeling regulation (21 CFR Part 801), please contact the Division of Industry and Consumer Education at its toll-free number (800) 638-2041 or (301) 796-7100 or at its Internet address

http://www.fda.gov/MedicalDevices/ResourcesforYou/Industry/default.htm. Also, please note the regulation entitled, "Misbranding by reference to premarket notification" (21CFR Part 807.97). For questions regarding the reporting of adverse events under the MDR regulation (21 CFR Part 803), please go to

http://www.fda.gov/MedicalDevices/Safety/ReportaProblem/default.htm for the CDRH's Office of Surveillance and Biometrics/Division of Postmarket Surveillance.

You may obtain other general information on your responsibilities under the Act from the Division of Industry and Consumer Education at its toll-free number (800) 638-2041 or (301) 796-7100 or at its Internet address

http://www.fda.gov/MedicalDevices/ResourcesforYou/Industry/default.htm.

Sincerely yours,

Mark N. Melkerson -S

Mark N. Melkerson Director Division of Orthopedic Devices Office of Device Evaluation Center for Devices and Radiological Health

Enclosure

# DEPARTMENT OF HEALTH AND HUMAN SERVICES Food and Drug Administration

# **Indications for Use**

Form Approved: OMB No. 0910-0120 Expiration Date: January 31, 2017

See PRA Statement below.

| 510(k) Number (if known)   | K150362   |
|--|---|
| K150362  | Page 1 of 1   |
| Device Name<br>NuVasive® CoRoent® Small Interbody System   |   |
| Indications for Use (Describe) The NuVasive CoRoent Small Interbody System is indicated for in mature patients. The CoRoent Small Interbody System is intended with cervical disc disease (DDD) at up to two contiguous levels fro supplemental fixation; the CoRoent SHL interbody device is require of supplemental fixation. The System is intended for use with autograncellous, cortical, and/or corticocancellous bone graft to facilitate who have had at least six weeks of non-operative treatment. | for use for anterior cervical interbody fusion in patients om C2 - T1. The System is intended to be used with red to be used with an anterior cervical plate as the form genous and/or allogeneic bone graft comprised of |
|  |   |
| Type of Use (Select one or both, as applicable)  |   |
| Prescription Use (Part 21 CFR 801 Subpart D)   | Over-The-Counter Use (21 CFR 801 Subpart C)   |
| PLEASE DO NOT WRITE BELOW THIS LINE – CONT   | INUE ON A SEPARATE PAGE IF NEEDED.  |
| FOR FDA USE (  | ONLY  |
| Concurrence of Center for Devices and Radiological Health (CDRH) (Sign   | ature)  |
| This section applies only to requirements of the *DO NOT SEND YOUR COMPLETED FORM TO THE   | •   |

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# 510(k) Summary

In accordance with Title 21 of the Code of Federal Regulations, Part 807, and in particular 21 CFR §807.92, the following summary of information is provided:

## A. Submitted by:

Martin Yahiro, M.D. Director, Medical Affairs NuVasive, Incorporated 7475 Lusk Blvd. San Diego, California 92121 Telephone: (858) 909-3360

Date Prepared: June 10, 2015

#### **B.** Device Name

Trade or Proprietary Name: NuVasive® CoRoent® Small Interbody System

Common or Usual Name: Intervertebral Body Fusion Device

Classification Name: Intervertebral Body Fusion Device with Bone Graft,

Cervical

Device Class II

Classification: 21 CFR § 888.3080

Product Code: ODP

#### **C.** Predicate Devices

The subject *CoRoent Small Interbody System* is substantially equivalent to multiple predicate devices. *NuVasive CoRoent Small Interbody System* (K140003) serves as the primary predicate device; while *NuVasive CoRoent System* (K081611), *NuVasive CoRoent Small Interbody System* (K140921), *NuVasive CoRoent Small Contoured Interbody System* (K142050), *Valeo*<sup>TM</sup> *Spacer System and Valeo*<sup>TM</sup> *II Interbody Fusion Device System* (K142264) are additional predicates.

## D. Device Description

The *NuVasive CoRoent Small Interbody System* is a hollow interbody cage manufactured from PEEK-Optima<sup>®</sup> LT-1 conforming to ASTM F2026. The implant contains a hollow core or graft aperture which allows for packing of autograft or allograft to help promote a solid fusion. Rows of teeth on the surface of each end of the device serve to grip the adjacent vertebrae to resist migration and expulsion of the device. The device includes marker pins composed of titanium alloy conforming to ASTM F136 and ISO 5832-3 or ASTM F1472 or tantalum conforming to ASTM F560 or ISO 13782. The pins serve as radiopaque markers allowing the location and orientation of the device to be seen radiographically during and after the procedure for position confirmation.

The implants are available in flat or contoured endplates, and come in a variety of different shapes and sizes to suit the individual pathology and anatomical conditions of the patient.



The subject device will be packaged and initially provided non-sterile, and is designed to be sterilized by the user before each use.

#### E. Indications for Use

The NuVasive CoRoent Small Interbody System is indicated for intervertebral body fusion of the spine in skeletally mature patients. The CoRoent Small Interbody System is intended for use for anterior cervical interbody fusion in patients with cervical disc disease (DDD) at up to two contiguous levels from C2 - T1. The System is intended to be used with supplemental fixation; the CoRoent SHL interbody device is required to be used with an anterior cervical plate as the form of supplemental fixation. The System is intended for use with autogenous and/or allogeneic bone graft comprised of cancellous, cortical, and/or corticocancellous bone graft to facilitate fusion. The cervical devices are to be used in patients who have had at least six weeks of non-operative treatment.

# F. Technological Characteristics

The subject *CoRoent Small Interbody System* is substantially equivalent to other predicate devices cleared by the FDA for commercial distribution in the United States. The subject device was shown to have equivalent technological characteristics to its predicate devices through comparison in areas including design, intended use, material composition, and function. This device does not contain software or electrical equipment.

#### G. Performance Data

The purpose of this 510(k) is to modify the Indications for Use for the subject *CoRoent Small Interbody System*. A clinical literature review was performed to support the use of the subject device at two levels. Based on the published clinical literature review, it was determined that the *CoRoent Small Interbody System* used in the treatment of two-level cervical degenerative disc disease has a safety and effectiveness profile that is similar to the predicate device. No other changes have been made to the interbodies since their clearance in the *CoRoent Small Interbody System* 510(k) K140003, *CoRoent Small Interbody System* 510(k) K140921, and *CoRoent Small Contoured Interbody System* 510(k) K142050. Therefore, no new nonclinical testing was performed for the purpose of this submission.

## H. Conclusions

Based on the indications for use, technological characteristics, clinical literature analysis, and comparison to predicate devices, the subject *CoRoent Small Interbody System* has been shown to be substantially equivalent to legally marketed predicate devices for its intended use.